(Atty. Docket No.: NEU-102.1P US)

Amendments to the Claims

This listing of claims will replace all prior versions and listings of claims in the application. Please amend the claims as indicated in the following Listing of the Claims, under the provisions of 37 C.F.R. § 1.121:

Listing of Claims

1. (currently amended) A method of treating an impaired neurological function in an individual who has sustained a brain injury comprising administering to said individual an effective amount of modafinil (benzhydrylsulfinylacetamide) in conjunction with a neurorehabilitation program comprising one or more neurostimuli designed to enhance or restore said impaired neurological function.

2. (canceled)

- 3. (currently amended) The method according to Claim 1 [[2]], wherein said neurorehabilitation program is selected from the group consisting of provides physical therapy, occupational therapy, speech therapy, and combinations thereof.
- 4. (currently amended) The method according to Claim <u>1</u> [[2]], wherein said neurorehabilitation program is selected from the group consisting of a physical/sensory protocol, an electrical and/or magnetic stimulation regimen, and/or a drug-based stimulation regimen.
- 5. (original) The method according to Claim 4, wherein said physical/sensory protocol comprises a neurostimulus selected from the group consisting of an exercise or task for motor function, an exercise or task for cognitive function, an exercise or task for a combination of motor and cognitive function, a light stimulation, an audio stimulation, a visual stimulation, a tactile stimulation, and combinations thereof.
- 6. (original) The method according to Claim 4, wherein said electrical and/or magnetic stimulation comprises trans-cranial magnetic stimulation (TMS) or deep brain stimulation (DBS)
- 7. (original) The method according to Claim 4, wherein said drug-based stimulation regimen comprises administering a neurostimulant drug.

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8. (original) The method according to Claim 7, wherein said neurostimulant drug is selected from the group consisting of caffeine, an amphetamine, a dextroamphetamine, a methylphenidate, and

combinations thereof.

9. (currently amended) The method according to Claim 1 [[2]], wherein said modafinil is administered to

said individual prior to or concurrently with said individual performing an exercise or task to promote or

restore an impaired neurological function.

10. (original) The method according to Claim 9, wherein administration of modafinil is stopped after

said individual performs an exercise or task and wherein said administration is not resumed until further

exercise or task is performed.

11. (currently amended) The method according to Claim $\underline{1}$ [[2]], wherein said administration of

modafinil and a neurorehabilitation program are ended after a period time, the individual is permitted a

period of rest from administration of said modafinil and said neurorehabilitation program, and the

individual is then administered modafinil and a neurorehabilitation program for a period of time.

12. (original) The method according to Claim 11, wherein after said period of rest, said administration

of modafinil is resumed at a different dose and/or said neurorehabilitation program is different from those

employed initially.

13. (original) The method according to Claim 11, wherein said period of administration of modafinil

and a neurorehabilitation program is 2 weeks and said period of rest is 4 to 12 weeks.

14. (original) The method according to Claim 1, wherein said modafinil is administered at a dose of

from 50 to 600 mg/day.

15. (original) The method according to Claim 14, wherein said dose is 100, 200, 400, or 600 mg/day.

16. (original) The method according to Claim15, wherein said dose is 200 mg/day.

17. (canceled)

18. (canceled)

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19. (canceled)

20. (original) The method according to Claim 1, further comprising administering to said individual a

dopaminergic agent that crosses the blood-brain barrier.

21. (original) The method according to Claim 1, further comprising administering to said individual a

dopaminergic agent selected from the group consisting apomorphine, bromocriptine, amantadine,

pergolide, pramipexole, ropinirole, fenoldopam, cabergoline, rotigotine, lysuride, talipexale, 7-OH DPAT,

quinpirole, SKF-38393, L-dopa, and combinations thereof.

22. (original) The method according to Claim 1, wherein the neurological function impaired in said

individual is selected from the group consisting of a cognitive function, a motor function, and a

combination of cognitive and motor functions.

23. (original) The method according to Claim 1, wherein said brain injury is the result of an event

selected from the group consisting of traumatic brain injury, an ischemic episode, spinal cord injury,

major organ failure, a brain injury associated with cardiovascular bypass surgery, an anoxic event, a

hypoxic event, a drug-induced brain injury, encephalitis, multiple sclerosis, and a degenerative disease.

24. (original) The method according to Claim 23, wherein said traumatic brain injury is the result of a

fall on a hard surface, a vehicle accident, or a strike to the head.

25. (original) The method according to Claim 23, wherein said ischemic event is a stroke.

26. (canceled)

27. (canceled)

28. (canceled)

29. (canceled)

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